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Site-specific dual-color labeling of long RNAs for single-molecule spectroscopy

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ABSTRACT

Labeling of long RNA molecules in a site-specific yet generally applicable manner is integral to many spectroscopic applications. Here we present a novel covalent labeling approach that is site-specific and scalable to long intricately folded RNAs. In this approach, a custom-designed DNA strand that hybridizes to the RNA guides a reactive group to target a preselected adenine residue. The functionalized nucleotide along with the concomitantly oxidized 3'-terminus can subsequently be conjugated to two different fluorophores via bio-orthogonal chemistry. We validate this modular labeling platform using a regulatory RNA of 275 nucleotides, the *btuB* riboswitch of *Escherichia coli*, demonstrate its general applicability by modifying a base within a duplex, and show its site-selectivity in targeting a pair of adjacent adenines. Native folding and function of the RNA is confirmed on the single-molecule level by using FRET as a sensor to visualize and characterize the conformational equilibrium of the riboswitch upon binding of its cofactor adenosylcobalamin. The presented labeling strategy overcomes size and site constraints that have hampered routine production of labeled RNA that are beyond 200 nt in length.

INTRODUCTION

Single-molecule (sm) spectroscopy along with Förster resonance energy transfer (FRET) has emerged as a versatile tool for probing distance distributions in biomacromolecules on the nanometer scale (1,2). Single-molecule detection can unravel conformational, kinetic, and thermodynamic heterogeneities of biomolecules like DNA, RNA, and proteins to provide information that is lost in ensemble experiments (3–6). An integral part of fluorescence spectroscopy is the choice and incorporation of suitable extrinsic fluorophores. smFRET requires a set of two spectrally

overlapping dyes, where the donor transfers its energy to the acceptor via dipole-dipole coupling. Working with long nucleic acids on the single-molecule level still remains challenging due to the lack of a universally applicable method that allows the introduction of a FRET dye pair in an orthogonal and site-specific manner while preserving the correct fold and optimal function of the biomolecule.

Many current approaches utilize modified nucleotides that are incorporated either chemically or enzymatically into the nucleic acid sequence for post-synthetic labeling. Most of these methods have either limitations concerning the size of the nucleic acid or lack site-specificity (7–9). Only recently, a platform for position-selective labeling of RNA has been introduced that combines T7 RNA polymerase transcription with solid-phase synthesis (SPS) (10). While being flexible in the type of labels, sequence upscaling is still problematic as yields are gradually decreasing with the number of repeating cycles. Moreover, ensuring selectivity within patches of consecutive identical nucleotides remains challenging. Currently, labeled RNAs of sizes smaller than 200 nt are usually prepared from custom-designed fragments which are ligated post-synthetically using T4 RNA or DNA ligase (11). Ligase-assisted approaches are, however, inherently restricted in the number and location of ligation sites. Intricate secondary structures complicate the design of the fragments to be ligated. In addition, production of the segments containing functionalized nucleotides still relies on solid phase synthesis and is therefore subject to size limitations. One labeling strategy which is independent of RNA size consists in hybridizing fluorophore that carry complementary sequences (DNA or PNA) to an *in vitro* transcribed RNA (12–14). However, this strategy often requires the insertion of non-native loops that may potentially alter the folding and function of the RNA. In case of the B₁₂ riboswitch dealt with herein, numerous attempts to introduce such inserts have led to RNA misfolding eventually rendering the switch inactive (data not shown). Ultimately, some methods use naturally occurring or *in vitro* selected enzymes to modify specific RNA substrates or in-

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sert a labeled mono- or polynucleotide at the 2'-ribose of an existing RNA strand (15–17).

Building on our recently developed strategy for DNA-templated generation of etheno-adducts within short, single stranded DNA (18) and RNA (19), we describe here a novel post-transcriptional and covalent labeling procedure for long RNAs with the ability to specifically graft an alkyne group on adenine residues buried in internal double-stranded regions (Figure 1A). In order to increase the accessibility of the target sites, complementary DNA oligonucleotides, known as helper sequences (dHSs) (20), are employed to disrupt the local secondary structure of the target RNA. During activation of the reactive strand (dRS) precursor, a concurrent oxidative opening of the 3'-terminal ribose ring to a dialdehyde moiety occurs and offers the possibility to place a second fluorophore at this position. The distinct orthogonal chemistry of the two functional groups at the internal and the 3'-terminal site allows for a site-specific dual-color labeling of RNA without the specter of cross-reactivity.

Here, we target a non-coding 275 nt long riboswitch that is embedded in the 5'-untranslated region (UTR) of the *btuB* gene from *Escherichia coli* (21,22). This B₁₂ responsive riboswitch regulates the expression of an outer-membrane transport protein that shuffles corrinoids like adenosylcobalamin (Ado-Cbl) into the periplasmic space (23,24). The RNA dynamics are monitored on the single-molecule level. The presented covalent and site-specific labeling protocol is novel in terms of RNA size (>200 nt) and its flexibility in choosing the most informative labeling sites while leaving the original sequence untouched.

MATERIALS AND METHODS

dHS assisted hybridization of a dRS analog

A mixture of *btuB* riboswitch (1 μ l, 1 μ M) and dHS (1 μ l, 0.1–50 μ M) buffered in 1 M sodium acetate, pH 5.5, was heated to 70°C for 5 min followed by incubation at 25°C for 30 min. Upon addition of the ³²P-labeled dRS analog (1 μ l, 1 μ M) the sample was put on ice for 10 min before loading it on a 6% (w/v) native polyacrylamide gel. The gel was run at room temperature, dried *in vacuo* and transferred onto a phosphor screen for scanning.

Site-specific modification of the *btuB* riboswitch

The *btuB* riboswitch (80 μ l, 36 μ M) buffered in 1 M sodium acetate, pH 5.5, was annealed (70°C, 5 min) with the respective dHS (1 eq. to *btuB* riboswitch). After incubation (25°C, 30 min) the freshly prepared dRS precursor (25 μ l, in 1 M sodium acetate, pH 5.5) was added and the solution was placed on ice for 10 min. Activation with sodium periodate (15 μ l, 20 mM, in 20 mM sodium acetate, pH 5.5) was carried out under mild agitation (25°C, 500 rpm, 90 min). The reaction was quenched by addition of ethylene glycol (30 μ l, 15 mM, in 20 mM sodium acetate, pH 5.5) and continued to incubate overnight. Desalting over a NAP-5 column, lyophilization and purification with 5% (w/v) denaturing PAGE afforded the functionalized *btuB* riboswitch devoid of dHS and dRS. The RNA was recovered from the gel by soaking the crushed pieces in a buffer

containing 3-(*N*-morpholino)propanesulfonic acid (MOPS, 10 mM, pH 6), EDTA (1 mM) and NaCl (250 mM), followed by EtOH precipitation. The internal alkyne modification was detected based on its characteristic fluorescence emission profile with a maximum around 415 nm upon excitation at 275 nm. The presence of the 3'-dialdehyde was demonstrated indirectly via coupling to the Cy3-hydrazide dye (*vide infra*).

Coupling of carbocyanine dyes to the modified *btuB* riboswitch

In order to label the 3'-terminal dialdehyde, the modified *btuB* riboswitch (25 μ l, 25 μ M, in 0.5 M triethylammonium acetate (TEAA), pH 7) was mixed with formamide (5 μ l, *purum*) and DMSO (5 μ l, *purum*), flushed with argon for 1 min and combined with argon-flushed Cy3-hydrazide (2 μ l, 10 mM in DMSO). Following incubation in the dark (25°C, 500 rpm, overnight), the RNA was precipitated with EtOH (75%). Subsequently, the sample was conjugated with sCy5-azide (sulfonated, Lumiprobe) without further purification to generate the double-labeled *btuB* riboswitch (*vide infra*).

The coupling of an azide-functionalized carbocyanine dye sCy5 (or sCy3, Figure 2) to the internal alkyne group was performed via a copper-catalyzed [3+2] cycloaddition (CuAAC). The 3'-modified *btuB* riboswitch (25 μ l, 25 μ M) was buffered with 0.5 M TEAA, pH 7, treated with formamide (10 μ l, *purum*), DMSO (55 μ l, *purum*), sCy5-azide (2 μ l, 10 mM in DMSO) and ascorbic acid (10 μ l, 5 mM) and flushed with argon for 1 min. A solution containing CuSO₄ and tris(benzyltriazolylmethyl)amine (TBTA) in a 1:1 ratio (5 μ l, 10 mM in 55% (v/v) DMSO) was added, followed by argon-flushing (1 min) and incubation in the dark (25°C, 500 rpm, overnight). Excess free dyes were removed by precipitating the RNA in EtOH, desalting the redissolved RNA over a NAP-5 column and purifying the crude product by 5% denaturing PAGE. Labeling yields ψ were calculated by comparing the absorption maxima at 260 nm (RNA) and 550 nm (Cy3) or 650 nm (sCy5) weighted by the respective extinction coefficient

$$\psi = \frac{A_{550 \text{ or } 650} \varepsilon_{btuB}}{A_{260} \varepsilon_{Cy3 \text{ or } sCy5}} \quad (1)$$

with $\varepsilon_{btuB} = 3022 \text{ mM}^{-1} \text{ cm}^{-1}$ (25), $\varepsilon_{Cy3} = 150 \text{ mM}^{-1} \text{ cm}^{-1}$ and $\varepsilon_{sCy5} = 250 \text{ mM}^{-1} \text{ cm}^{-1}$ (26).

Additional methodological information is available in the supporting information.

RESULTS AND DISCUSSION

In the following, we will outline the steps involved in the design of suitable reactive and helper strands, their hybridization to the riboswitch and subsequent coupling to the fluorophores. We further evaluate the site-specificity of the modification and prove the integrity of both the RNA and the dyes upon bioconjugation via fluorescence spectroscopy.

Selection of the labeling sites

The *btuB* riboswitch construct comprises the full-length aptamer and expression platform (240 nt) and is extended

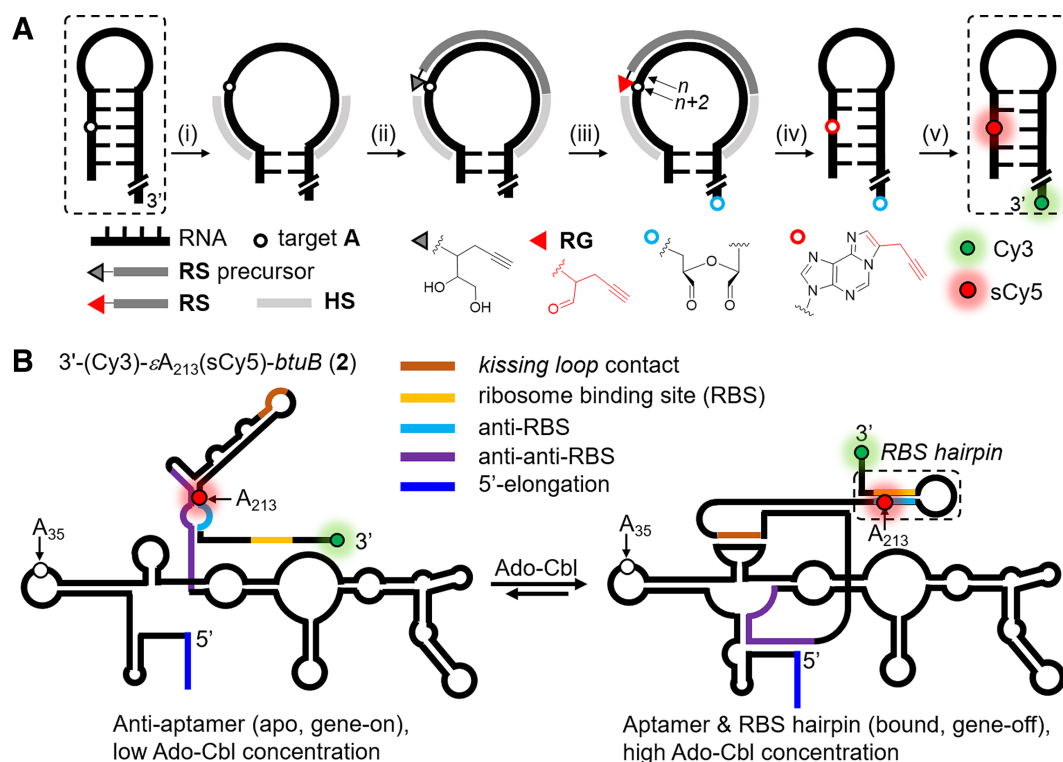


Figure 1. Site-specific dual-color labeling of long RNAs. (A) Schematic illustration of the double labeling workflow at an internal adenine and the 3'-end. (i) dHS annealing, (ii) hybridization of the dRS precursor, (iii) activation by sodium periodate, (iv) transfer of the reactive group and modification of an internal adenine, (v) orthogonal fluorophore coupling with Cy3-hydrazide at the 3'-end dialdehyde and sCy5-azide at the internal alkyne-εA via CuAAC. (B) Depiction of the translational control by the *btuB* riboswitch (22): the equilibrium between two conformers is shifted in response to the cofactor adenosylcobalamin (Ado-Cbl). In the apo form, the Shine-Dalgarno sequence (or ribosomal binding site, RBS) is accessible to the ribosome. Conversely, when Ado-Cbl levels increase, the RNA establishes a kissing loop interaction whereupon the RBS is sequestered through formation of an RBS hairpin motif. This equilibrium can be probed via single-molecule FRET by placing a donor fluorophore (Cy3) at the 3'-terminus and an acceptor (sCy5) at A₂₁₃ or A₃₅ (indicated with arrows).

at the 5'-end by 35 nucleotides of the natural genomic sequence to allow surface immobilization for total internal reflection fluorescence (TIRF) microscopy (Figure 1B and Supplementary Figure S1). Adenines 35 (A₃₅, note that the 5'-elongation is not part of the numbering scheme) and adenine 213 (A₂₁₃) are chosen as the labeling sites for either one of the sulfo-carbocyanine fluorophores sCy5 or sCy3. A₃₅ is located in an internal hairpin loop in close proximity to the kissing loop that forms upon binding of the cofactor Ado-Cbl (Figure 1B). The expression platform harbors a second modification site, A₂₁₃, which is particularly challenging to target as it is situated in a duplex region. In conjunction with a suitable fluorophore at the 3'-end, the FRET intensity reflects the underlying conformational equilibrium that characterizes the sequestration of the ribosomal binding site (RBS) during translational control by the riboswitch.

The synthesis of the reactive group (RG), as part of the reactive strand (dRS) was described previously (Supplementary Figures S2 and S3) (19). No adenines and cytosines are placed next to the RG in the dRS in order to prevent self-modification (Supplementary Figures S6 and S7) (18,19). For reasons of clarity, we refer to the target ribonucleotide opposite the one linked to the RG as being the *n* position (Figure 1A). The RNA residue which is next to *n* but not hydrogen-bonded to the dRS is in the *n* + 1 position. Both dRSs were designed such that the target adenines are sit-

uated in the *n* + 2 position, which is the ideal position due to the linker length of the RG (Supplementary Figure S3) and fulfills the criterion that the dRS lacks a terminal A or C base. We previously demonstrated that labeling efficiencies for the *n* + 1 and *n* + 2 positions are similar and then progressively diminish with increasing distance from the RG (18,19). This makes designing the dRS more flexible since more labeling sites are tolerated.

Helper sequences assist dRS annealing

Targeted delivery of a functional group to a nucleic acid by DNA-templated synthesis (27) has so far only been applied to single-stranded DNA (18) or long single-stranded regions of RNA molecules (19). The complex secondary and tertiary architecture of the folded *btuB* riboswitch restricts the accessibility of certain ribonucleotides and hampers hybridization of the dRS to the RNA. To protect the integrity of both the RNA and the RG, we avoid exposure to high temperatures during the modification process and melt the local base-pairing regime by annealing complementary dHSs (15,20) at moderate temperatures up- or downstream of the RNA sequence complementary to the dRS before activating the RG *in situ* (Figure 1A). The dHS is designed in such a way to sufficiently disrupt the secondary structure elements in proximity of the desired labeling site. The length

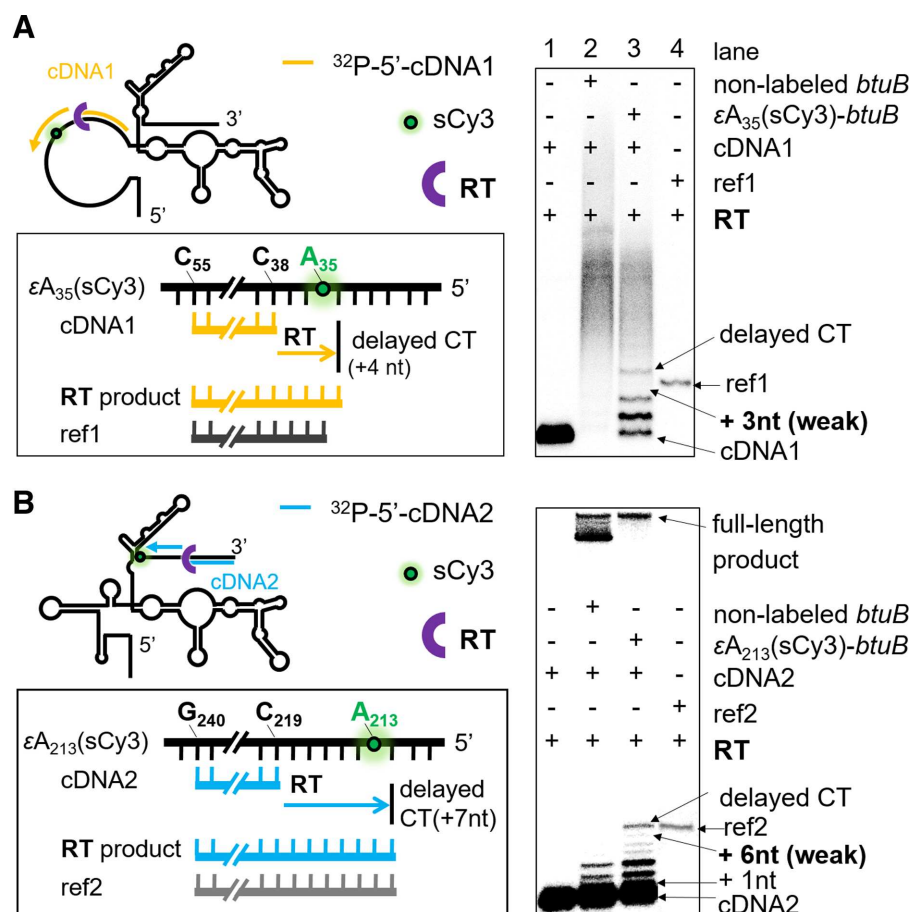


Figure 2. Evaluating the site-specificity of internal labeling by reverse transcription. Internally labeled *btuB* riboswitch constructs $\epsilon A_{35}(sCy3)$ -*btuB* (A) and $\epsilon A_{213}(sCy3)$ -*btuB* (B) are used as templates for the reverse transcription of short cDNA strands. The autoradiograph of the denaturing, polyacrylamide footprinting gel (18% w/v, 7 M urea) shows the original cDNA (lane 1) and the extended cDNA in presence of the non-labeled *btuB* riboswitch (lane 2) or the internally labeled *btuB* riboswitch (lane 3). Short DNA oligonucleotides, cDNA1 + 3nt (A) and cDNA2 + 7nt (B) are shown as references (lane 4). The weak band at cDNA + 3 nt (A) or cDNA + 6 nt (B) along with the stronger band at cDNA + 4nt (A) and cDNA + 7nt (B) reflect a delayed chain termination (CT) upon insertion of one additional nucleotide past the modification site, as previously observed (31).

of the **dHS** majorly depends on the RNA secondary structure and the GC content and accordingly on the predicted melting temperature.

In case of A₃₅, two helper sequences (**dHS1a** and **dHS1b**) are designed to mask short segments on either side of the target nucleotide (Supplementary Figure S4A) and thereby open up the stem loop temporarily. To assess the efficiency of the **dHS** in promoting **dRS** hybridization, a ³²P-labeled DNA oligonucleotide, referred to as **dRS** analog, was added to the *btuB* RNA in presence of increasing amounts of **dHS**. In the absence of any **dHS** only 60% of the **dRS1** analog bind to the RNA at room temperature. In contrast, treatment with one equivalent of **dHS1a** and **dHS1b** each increases the binding efficiency of **dRS1** to >95% at ambient temperature (Supplementary Figure S4B and C). By comparison, temperature induced unfolding shows only ~70% hybridization efficiency in absence of any **dHS**. More evidently in case of A₂₁₃, only about 20% of the **dRS2** analog anneal to the RNA in the absence of **dHS2** at room temperature, whereas addition of one equivalent of **dHS2** leads to ~75% bound **dRS2** (Supplementary Figure S5b–c). Again, thermal melting without **dHS2** produces no more than

50% annealed DNA. In brief, the **dHSs** enable efficient hybridization of the **dRS** at room temperature – a prerequisite for an accurate positioning of the **RG** close to the target sites A₃₅ and A₂₁₃ where the functional alkyne group should be transferred onto.

RNA functionalization in a one-pot reaction

After annealing of the **dHS** and the **dRS** to the RNA, activation of the **RG** *in situ* under mild oxidative conditions to a reactive aldehyde is achieved by sodium periodate (Supplementary Figure S6–S7). At the same time, the two hydroxyl groups of the 3'-terminal ribose unit are oxidized to a dialdehyde (28,29). At the internal site, the exocyclic amine of adenine forms an etheno adduct (ϵA) with an additional alkyne moiety. Etheno bridge formation is detected by exciting the RNA at either 275 nm or 308 nm to produce an emission peak around 415 nm that is characteristic of ϵA (Supplementary Figure S8) (30). As a result of this one-pot reaction the riboswitch is modified with two chemically orthogonal functional groups, one at an internal adenine and another at the 3'-terminus. Fluorophore coupling can now be achieved by treating the functionalized RNA with azide-

and hydrazide-functionalized fluorophores without worrying about dye cross-reactivity.

Site-specificity of the internal labeling scheme

As stated previously, the site-specificity of the internal nucleotide modification is crucial for subsequent single-molecule experiments. A polymerase stop assay was applied (18,19), in which a cDNA primer anneals downstream of the putative labeling site and is extended by a reverse transcriptase (RT). Two dye labeled constructs, $\varepsilon A_{35}(\text{sCy3})\text{-}btuB$ and $\varepsilon A_{213}(\text{sCy3})\text{-}btuB$, are used to assess site-specificity of the internal labeling. The footprinting pattern (Figure 2a) around fluorophore position A_{35} shows distinct bands corresponding to the cDNA primer extended by 1, 2 and 4 nt. The RT is not stalled at the presumed labeling site (cDNA1 + 3 nt) but inserts one additional nucleotide before stopping and releasing the nascent cDNA, a process also known as delayed chain termination (CT) (31). For εA_{213} (Figure 2b), several bands representing cDNA2 + 1–3 nt are observed for both the non-labeled riboswitch and the $\varepsilon A_{213}(\text{sCy3})\text{-}btuB$ construct, which reflects the stalling of the RT during initiation of primer extension (32). The additional bands in the vicinity of the labeling site (cDNA2 + 6 nt) are very weak. Major pausing of the RT occurs again one nucleotide after the modification site. In combination with the fluorescence emission profile of εA (Supplementary Figure S8), this is a clear indication that etheno bridge formation occurs predominantly at A_{35} and A_{213} , whereas εA_{36} , and εC_{12} are present in trace amounts only, if at all. Moreover, labels at those sites would only slightly affect smFRET measurements as mono-labeling at $n \pm 1$ or 2 nt changes the transfer efficiency by about 0.08 in the worst case (calculated for a helical rise of 0.28 nm with maximal sensitivity around the Förster radius $R_0 = 5.4$ nm). Importantly, and in contrast to current ligation approaches, our labeling strategy prohibits non-templated nucleotide incorporation and thus the underlying RNA sequence remains unchanged.

Double labeling yields are sufficient for single-molecule experiments

The reaction of 3'-terminal periodate-activated sugars with hydrazide functionalized fluorophores proceeds with relatively high yields (29). To avoid reduction and hence inactivation of the 3'-dialdehyde during the double-labeling procedure the functionalized *btuB* riboswitch was first incubated with a hydrazide derivatized Cy3-dye. Only subsequently, CuAAC Click chemistry under reductive conditions was used to couple the azide derivative of sCy5 to the alkyne modified internal adenines A_{35} or A_{213} , respectively (Supplementary Figure S9). Co-localized green and red emission on denaturing gels (Figure 3A and Supplementary Figure S12A) along with the slightly red-shifted absorption and fluorescence emission spectra, as compared to the free dyes (Supplementary Figure S10), indicates that both fluorophores are covalently attached to the riboswitch constructs 3'-(Cy3)- $\varepsilon A_{35}(\text{sCy5})\text{-}btuB$ (1) and 3'-(Cy3)- $\varepsilon A_{213}(\text{sCy5})\text{-}btuB$ (2). Labeling efficiencies (Eq. 1) are 26% (Cy3) and 14% (sCy5) for construct 1 and 37% (Cy3) and 17% (sCy5) for construct 2, respectively. Double

labeling yields are about 5% which is generally sufficient for single-molecule fluorescence spectroscopy. In comparison, classical SPS of RNA sequences proceeds with ~98% yield per nucleotide, affording an overall yield of ~0.4% for an RNA of 275 nt in length. This yield will be further reduced by post-synthetic labeling. In our case, the final yields result from four independent reactions: (i) internal adenine modification, (ii) 3'-end modification, (iii) internal dye coupling via CuAAC reaction, and (iv) 3'-end dye coupling. Step (i) in particular is sequence dependent and yields are expected to vary depending on the accessibility of the modification site. We purposely chose sites that are buried and thus not easily tractable, thereby making the use of HS sequences essential and pushing the method to its limits. Labeling of a 633 nt long RNA in single stranded regions has been shown previously (19). Here, we demonstrate that RNAs, irrespective of their length, can be double-labeled and not only in internal loops but also in helical duplex regions. Proper folding of the single- and dual-color-labeled RNA constructs was evaluated by fluorescent PAGE (Supplementary Figure S11). All constructs show the same migration behavior as the non-labeled one and hence adopt similar 3D folds. Accordingly, the dyes do not interfere with the folding of the RNA.

Photophysics of cyanine dyes within the riboswitch environment

Carbocyanine fluorophores are known to be sensitive to their local environment which is reflected in changes of the fluorescence lifetime, quantum yield and anisotropy (33–35). The torsional flexibility of the polymethine chain of carbocyanines causes an isomerization to a non-fluorescent mono-*cis* state. The photoisomerization reaction is an excited state process that is conditioned by sterical constraints imposed on the fluorophore. Interactions with the biomolecular environment define the relative populations of *trans* and *cis* states, which are experimentally accessible via a change in molecular brightness. In other words, the RNA shapes the photophysics of the fluorophore, a phenomenon referred to as RNA-induced fluorescence enhancement (RIFE) (34). In order to demonstrate the integrity of the dyes, we measured fluorescence lifetimes of the dyes prior to and after coupling to the riboswitch (Figure 3B and Supplementary Figure S12B). While the fluorescence of the free azide form of sCy5 decays monoexponentially with a lifetime of $\tau = 0.76 \pm 0.03$ ns, a second lifetime component ($\tau_2 = 1.9 \pm 0.1$ ns) emerges upon covalent attachment to εA_{213} (Figure 3B). Both components have a relative weight of roughly 50% (Supplementary Table S2). The lifetime decay of the 3'-terminal Cy3 can also be represented by a biexponential function ($\tau_1 = 0.67 \pm 0.02$ ns and $\tau_2 = 2.6 \pm 0.1$ ns). Compared to the free dye in solution ($\tau = 0.11 \pm 0.01$ ns) the lifetime is drastically increased, indicating a considerable reduction in torsional flexibility and photoisomerization as expected.

The extent of motional restriction by the RNA environment is illustrated by recording the fluorescence anisotropy as a function of time (Figure 3C and Supplementary Figure S12C). The decays are modeled in terms of a fast, hindered rotation (wobbling-in-cone model) governed by the

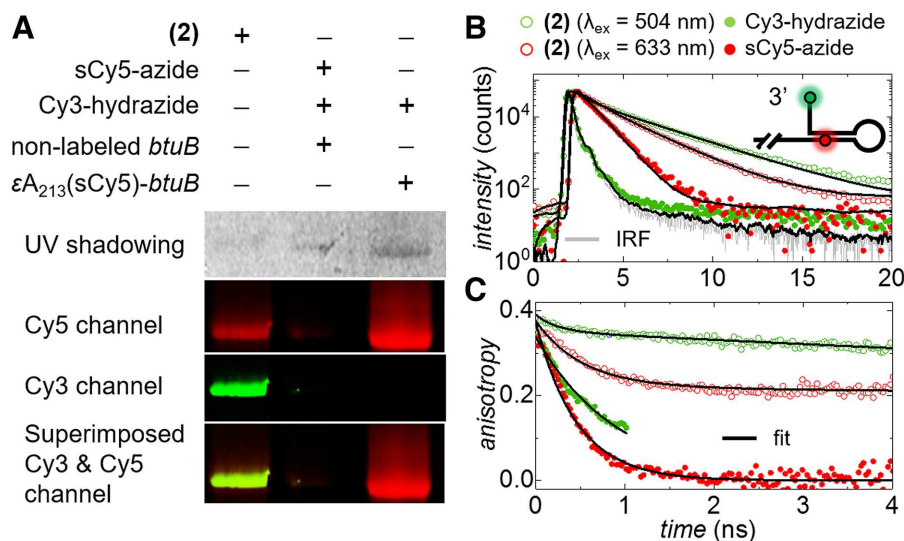


Figure 3. Photophysical characterization of the double-labeled *btuB* riboswitch. (A) Characterization of the construct 3'-(Cy3)- ϵA_{213} (sCy5)-*btuB* (2) by denaturing PAGE (10% w/v, 7 M urea). Bands on the gel are visualized by UV-shadowing ($\lambda_{ex} = 254$ nm, top) or fluorescence scanning (excitation at 532 nm or 635 nm, respectively, bottom). Color-coded channels show Cy3 (green) and Cy5 (red) emission. Co-localized dyes appear as yellow bands. Control lanes with the unmodified *btuB* in the presence of both free dyes and controls with the Cy5-labeled construct in the presence of free Cy3 show no fluorescence signal from the non-covalently attached dye and thus intercalation can be excluded. (B) Fluorescence lifetime decays of 2 excited at 504 nm (Cy3) and 633 nm (sCy5) compared to the free fluorophores in solution. The instrument response function (IRF) is drawn in gray. (C) The dynamic anisotropy reflects the orientational constraints imposed on the fluorophores by the RNA microenvironment.

fluorophore linker (τ_r^{local}) and a decoupled, global tumbling (τ_r^{global}) component along with the entire construct (Eq. S2) (34). The long global correlation time ($\tau_r^{\text{global}} > 40$ ns) and high residual anisotropy ($r_\infty > 0.22$) of the fluorophores suggest considerable stacking of the fluorophores on the ribonucleic acid backbone as observed previously for various RNA systems (Supplementary Table S2) (34). The non-sulfonated Cy3 is intrinsically more hydrophobic and displays a higher interaction propensity with the RNA. Moreover, it is directly coupled to the 3'-end which allows the dye to additionally stack onto the terminal guanine (36). The reduced local correlation time of the 3'-Cy3 ($\tau_r^{\text{local}} \sim 0.2$ ns) in comparison to Cy3 free in solution ($\tau_r^{\text{local}} = 0.92$ ns) as well as the high residual anisotropy ($r_\infty = 0.35$) reflect the dye wobbling within a small cone angle and support the discussion above. The internal modification with sCy5, however, shows about 50% free dye rotation and no reduction of the local correlation time. Even though the tumbling of the biomolecule-tethered fluorophores is spatially constrained, the dyes retain full functionality. It is noteworthy that local confinements on the fluorophore movement affect the orientation parameter κ^2 in the Förster radius R_0 and thus need to be considered for exact distance measurements.

Riboswitch dynamics monitored by smFRET

Single-molecule spectroscopy has the key advantage to dissect the ensemble and monitor individual molecules over time. Total internal reflection fluorescence (TIRF) microscopy usually requires the molecule of interest to be tethered to the surface of a quartz slide (37). For that purpose, a biotin-tagged DNA oligonucleotide is hybridized to the 5'-elongation of the riboswitch (Supplementary Figure S13a). Applying an alternating-laser excitation (ALEX)

(38) scheme with illumination at 532 and 640 nm, time traces of the dual-color-labeled RNA are recorded in the absence and presence of the metabolite Ado-Cbl. The latter is reported to quench the fluorescence emission of the Cy3 donor to some extent (4,23). Nonetheless, we are able to detect the molecules upon Ado-Cbl binding and observe the energy transfer. The 3'-(Cy3)- ϵA_{213} (sCy5)-*btuB* (2) construct probes the formation of the RBS hairpin (Figure 4A). Abrupt single-step photobleaching of Cy3 and sCy5 within the observation time along with a stable stoichiometry of Cy3 (S_{Cy3}) fluctuating around 0.5 suggests that both dyes are covalently attached to the same RNA molecule (Supplementary Figure S13B). The riboswitch interconverts between a low and a high FRET state in the presence of 3 mM Mg^{2+} (Figure 4B and Supplementary Figure S14) (39). Upon addition of the cofactor the relative populations of the states shift in favor of the high transfer efficiencies. FRET time traces are analyzed with a home-written Matlab software (40) to build cumulative histograms (Figure 4C). In addition to the high and low FRET states, we find a substantial contribution of intermediate FRET states originating from static molecules (Supplementary Figure S14, static molecule 2). Additionally, we observe dynamic molecules that show short excursions to either the high or the low FRET state (Figure 4b and Supplementary Figure S14, dynamic molecule 2 and 3). Interestingly, consecutive transitions from low to intermediate to high FRET and vice versa are barely visible. Intermediate states between 0.2 and 0.8 may arise from time-averaging (41), but can also be attributed to trapped conformations of the riboswitch expression platform. Further, shifting FRET states can arise from quantum yield changes of carbocyanine fluorophores due to differences in the photoisomerization probability in the local RNA environment, as reported

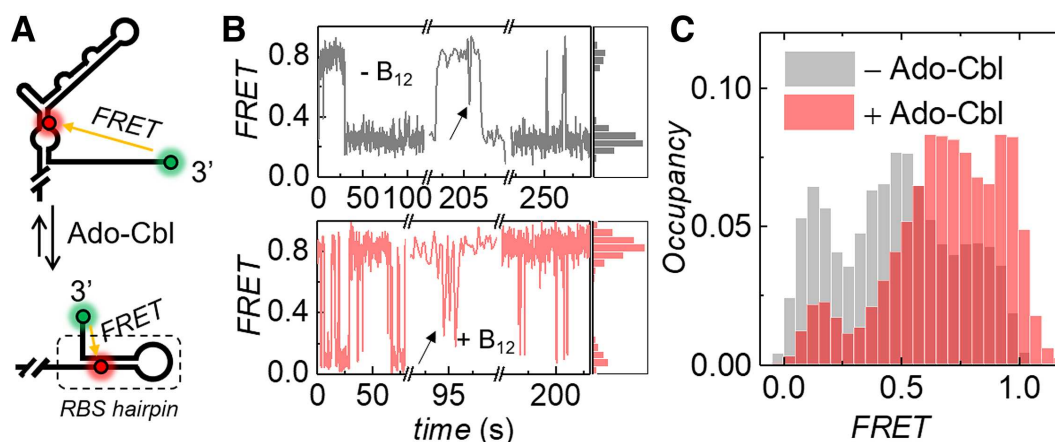


Figure 4. Single-molecule FRET characterization of 3'-Cy3- ϵ A₂₁₃(sCy5)-*btuB* (2). (A) Formation of the RBS hairpin upon addition of Ado-Cbl (B₁₂). (B) FRET trajectories in the absence and presence of 0.1 mM Ado-Cbl (50 mM KCl, 3 mM MgCl₂). FRET trajectories have been partly rescaled to visualize fast state transitions from/to intermediate states (indicated by black arrows). (C) Normalized cumulated FRET histograms built up from about 90 molecules showing a predominant occupancy of high FRET values in the presence of the metabolite (red).

recently (34,42). Indeed, Stoichiometry-FRET histograms show substantial broadening which supports an influence of dye photophysics (Supplementary Figure S13C). A more detailed analysis and characterization of the *btuB* folding pathway will be addressed elsewhere. Importantly, different population levels of lower and higher FRET states in response to Ado-Cbl provide evidence of a tunable equilibrium between the apo and the bound conformation as observed previously for a hydroxycobalamin riboswitch (4).

CONCLUSION

Current efforts in co- and post-transcriptional labeling of nucleic acids are directed towards finding the best trade-off between site-specificity, programmability and size of the target molecule while preserving the integrity of the native RNA structure and folding. While PLOR is a versatile platform adaptable to many different applications, the practicability for very long RNAs remains elusive. Ligase-assisted approaches on the other hand have been used to prepare riboswitch domains with up to 200 nt, but suffer from restrictions in the number and positioning of appropriate ligation sites. Non-covalent hybridization approaches are generally independent of the RNA dimensions, however they usually require changes made to the native sequence in order to accommodate the fluorescently labeled complement.

We developed a minimally invasive, nucleotide- and position-selective labeling strategy which is scalable to RNAs beyond 200 nt in length. Our method utilizes the selective annealing of custom-designed DNA probes to modify an *in vitro* transcribed RNA *in situ* in principle at any specific adenine or cytosine. Regions that feature buried nucleotides such as internal stem loops and duplex structures are tackled by introducing additional dHSSs, which work synergistically with the dRS to disrupt the local secondary structure. The DNA-templated transfer of a functional group to an adenine and subsequent coupling with a fluorescent probe is shown to preserve the overall RNA fold and function. Cross-contamination in dual-color labeling is avoided by using two different bio-orthogonal coupling re-

actions. The presented labeling platform has been validated on the ensemble and single-molecule level by means of TC-SPC and smFRET using a long riboswitch as a model system. The modular nature of the method allows the design of alternative functional handles including azides, amines, or thiols to pave the way for three-color FRET. On the RNA side, the scope of possible modification targets can be readily extended to cytosines which also feature an exocyclic amine group. Overall, this study provides a novel approach to fluorescently label long nucleic acids. It is highly customizable with respect to the location of the labeling sites and hence particularly suitable for studying large and sensitive RNA systems such as riboswitches or ribozymes.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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